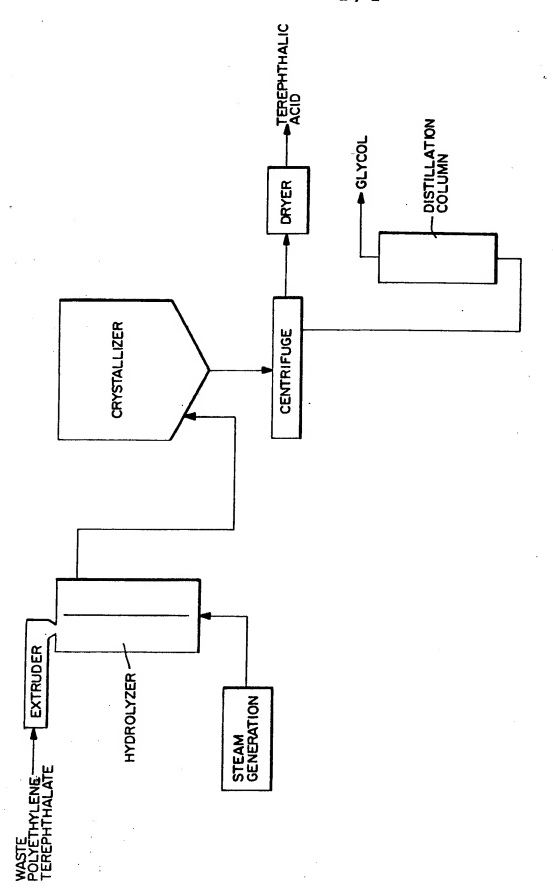
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- (54) Continuous procedure for obtaining pure terephthalic acid and glycol starting from polyethylene terephthalate waste
- (57) A procedure is described to obtain pure terephthalic acid and glycol, which consists in carrying out, in a continuous manner and in the presence of decolorizing carbon, an aqueous and neutral hydrolysis of polyethylene terephthalate waste, under high temperature and pressure conditions, continuously filtering the product at the hydrolysis temperature and crystallizing the terephthalic acid, while the mother liquors are subjected to distillation in order to obtain the glycol.

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	SPECIFICATION	
5	Continuous procedure for obtaining pure terephthalic acid and glycol starting from polyethylene terephthalate waste	5
	Field of the invention  The invention under consideration is related to a continuous method which permits obtaining pure terephthalic acid and glycol and more specifically, it is related with a procedure which utilizes as a raw material polyethylene terephthalate waste and in which it is not necessary to first separate the crude or impure terephthalic acid in order to purify it in a subsequent stage by means of one of the known procedures. In the method of the invention under consideration, the product is obtained pure and can be utilized as a raw material once again in the preparation of the polyethylene terephthalate utilized in the manufacture of polyester fibers and plastics.	10
15	Description of the prior art  Neutral hydrolysis of polyethylene terephthalate at a high temperature and pressure is a known operation (see Ludewig, H. and Ramm, H.: German Economic Patent 14,854 (1956) and Littmann, E. On the preparation of terephthalic acid or its dimethyl or diglycol ester from	15
20	polyethylene terephthalate. Abh. d. Deutsch, Akad. d. Wiss. Kl. F. Chem, Geolog. und Biochemie I (1963) 401–411).  Neither of the methods described in the aforementioned references mentions the addition of decolorizing carbon in the hydrolysis stage.	20
25	On the other hand, the methods mentioned there are discontinuous methods in which, moreover, the terephthalic acid is obtained impure so that once it is separated, it will have to be subjected to some purifying procedure.  In this way, the methods to which reference has been made require a subsequent purifying stage which may be the dissolution of the product in sodium hydroxide and subsequently, a	25
30	reprecipitation by means of acidification, either with treatment of the solution obtained or without an intermediate treatment, or by having recourse to transformation to the dimethyl ester, which is subsequently sublimed.	30
35	Summary of the invention  The method of the invention under consideration has as its object the obtaining of terephthalic acid and glycol in which the purification is undertaken simultaneously with the hydrolysis.  Another object of the invention under consideration is to provide a method for the preparation of pure terephthalic acid and glycol in which the number of stages of the process is reduced.  An advantage of the method of the invention under consideration is that upon reducing the number of stages of the process—that is, upon eliminating the subsequent stage of purifying	35
40	polyethylene terephthalate in a simple manner.  Yet another advantage of the method of this invention is that due to the simplicity of the	40
45	equipment utilized, it can be carried out with relative ease.  These and other objects of the invention under consideration will, in part, be obvious and, in part, will become more apparent as the description proceeds and with the aid of the detailed description of the invention.	45
50	Detailed description of the invention  According to the method of this invention, the polyethylene terephthalate waste is supplied in the form of granules or yarns or in any other form suitable for handling, to the melting chamber of a screw-type extruder so that the extruder, in turn, feeds it, in a continuous manner, to a hydrolyzer. Alternatively, the polymer can also be melted and injected continuously into the hydrolyzer in this state by means of a high-pressure pump.	50
55	In a simultaneous manner, water is injected into said hydrolyzer in a suitable porportion—that is, from 3 to 20 times the amount of the polymer. The water can be introduced cold or hot. In turn, the decolorizing carbon is added in a proportion between 0.1% and 30% (with respect to the quantity of the treated waste), feeding it to the hydrolyzer either with the polymer or suspended in water. Also, a large amount can be introduced in bulk intermittently into the	55

hydrolyzer every so often. The hydrolyzer utilized can be of any type—that is, it can be

The hydrolyzer, in turn, should be equipped with a source of heating which can be a jacket

60 cylindrical vertical, horizontal or inclined, or it can be the U-shaped type, either vertical,

horizontal or inclined.

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Heating can also be provided by means of live steam at high pressure, which is introduced directly into the hydrolyzer.

Preferably, the hydrolyzer should be equipped with devices which impart a slight agitation.

When the equipment is put into operation, the polymer which has not yet been hydrolyzed will be found in the lower part. This is also true for the greater part of the carbon, whereas, on the other hand, the supernatant liquid will be an aqueous solution of terephthalic acid and glycol with some proportion of the decolorizing carbon utilized, which will be suspended in the

This solution is subjected to a filtering operation under the temperature and pressure 10 conditions in which the hydrolysis is carried out, with the object of eliminating the carbon and all the impurities adsorbed by it. This filtration can be carried out in many diverse ways but the use of a heated cartridge filter is particularly preferred; this is located next to the hydrolyzerthat is, said filtration can be undertaken by means of cartridges which are installed directly within the hydrolyzer.

Once the solution is filtered, the pressure to which it is subjected is released and it is sent to a 15 15 continuous crystallizer which may be at atmospheric pressure, at a pressure higher than atmospheric pressure, or at a pressure lower than atmospheric pressure.

This crystallizer can be equipped with a cooling jacket or a coil, or cooling may be attained by simply venting the pressure.

20 The magma obtained in this manner in the crystallizer is filtered continuously, either by using 20 a filter or with a centrifuge. The crystals are washed in the same filtering device and are dried so as to be stored and recycled.

On the other hand, the filtrate of the magma, which is an aqueous glycol solution is sent to distillation equipment where the glycol is obtained in pure form.

The following indicates the preferred operating conditions in the method, with the upper and 25 lower limits of the permissible conditions being indicated. The indicated parameters are not independent of one another and upon modifying one of them, it may be necessary, or at least convenient, to change one or several of the others.

	Minimum	Maximum	Remarks	30
Water/polymer ratio	3	20	Note 1	
Carbon/polymer ratio	0.1%	30%	Note 2	
Hydrolysis temperature	200°C	300°C	Note 3	
Hydrolysis pressure	15 atmospheres	100 atmos- pheres	Note 4	35
Residence time in the hydrolyzer Residence time in the	5 min.	6 hours	Note 5	
	5 min.	5 hours	Note 6	40
Temperature in the crystallizer	– 10°C	200°C	Note 7	
	Carbon/polymer ratio Hydrolysis temperature Hydrolysis pressure  Residence time in the hydrolyzer Residence time in the crystallizer Temperature in the	Water/polymer ratio Carbon/polymer ratio Hydrolysis temperature Hydrolysis pressure  Residence time in the hydrolyzer Residence time in the crystallizer Temperature in the	Water/polymer ratio Carbon/polymer ratio Hydrolysis temperature Hydrolysis pressure  Residence time in the hydrolyzer Residence time in the crystallizer Temperature in the  3 20 30% 30% 300°C 100 atmospheres 100 atmospheres 6 hours 5 min. 5 hours	Water/polymer ratio 3 20 Note 1 Carbon/polymer ratio 0.1% 30% Note 2 Hydrolysis temperature 200°C 300°C Note 3 Hydrolysis pressure 15 atmospheres 100 atmospheres Residence time in the 5 min. 6 hours Note 5 hydrolyzer Residence time in the crystallizer 5 min. 5 hours Note 6 Temperature in the

Note 1. The quantity of water should be sufficient so as to keep the terephthalic acid dissolved as it is being formed; such a quantity will depend on the temperature at which the hydrolysis takes place.

Note 2. The quantity of carbon needed will depend on the quantity of impurities in the polymer and the final purity desired in the terephthalic acid produced.

Note 3. The greater the temperature, the higher will be the rate of the reaction, with the pressure in the equipment being greater at the same time.

Note 4. The pressure is a function of the temperature.

Note 5. The residence time required will depend on the temperature used and on the morphology of the polymer.

Note 6. The residence time in the crystallizer will depend on the average size of the crystal desired, with the crystal being larger, the greater the residence time.

In accordance with the data previously indicated, the method of the invention can be carried out under conditions which are different up to a certain point, and it can be modified in accordance with the level of production required.

Also, very different heating sources can be used, with the needed changes also made in 60 accordance with the morphology and the quality of the waste which is utilized as a raw material. There will also be variations in accordance with the characteristics of purity and size of the crystal desired in the terephthalic acid produced.

Taking into consideration all the modifications which are comprised within the spirit and scope of the invention, the following will give the preferred embodiment, which is merely for illustrative purposes.

	mustrative purposes.			
5	Detailed explanation of the preferred embodiment In this embodiment, the method of the invention is carried out with the following equipment as illustrated in the drawing:			
		is discharged by means of a multiple-orifice spray head		
10	b) A vertical cylindrical hydrolyzer which hydrolyzer into two semiconductor section length but rather will leave a free space by	ch has a vertical plate in its interior which divides the ns. Said plate does not have to extend along the whole both in the upper as well as the lower parts. The supply der is centered over one of the semicircles. On the	10	
15	upper part of the other semicircle there is connected with a common discharge tube the side corresponding to the supply of the equipped with two connections—one who	s a bundle of filter cartridges whose outlets are all e. The entry of water is on one side of the hydrolyzer, he polymer. The bottom of the hydrolyzer will be introduction of live steam, and	15	
20	equipped with an agitator. The crystallize cartridges on the hydrolyzer by means of	ade up of a vertical tank, a conical bottom and which is it is connected to the discharge tube of the filter a line and an automatic valve which permits the	20	
	in the hydrolyzer. Said crystallizer also in	small quantity of steam which has an agitation function cludes an automatic level control discharge device. ning device which receives the discharge from the		
25	e) A drief lotter and which, in turn, continuou  e) A drief of the vibrating screen type.	sly discharges towards the drier.  of the glycol, on the basis of mother liquors.	25	
	What has been described in the precede preferred for carrying out the method, when the precede preferred for carrying out the method.	ling is related to the equipment which is particularly		
30	The hydrolyzer is loaded with a suitable of a metering pump. High-pressure saturates of the hydrolyzer. The waste polyme	e quantity of water in a continuous manner by means ated steam is also injected continuously from the lower r, together with decolorizing carbon is fed continuously	30	
35	to the hydrolyzer by means of the extrud.  The discharge from the hydrolyzer is a means of the excess steam.	djusted in such a manner that it is slightly agitated by	35	
	Once the hydrolysis has been undertak	en, the product is subjected to filtration in order to be licate that the agitation in the crystallizer has to be the pended.		
40	The crystals obtained are subjected to the flow of air and the temperature suital	a drying stage, which is carried out in a drier which has	40	
	the glycol pure.	specific reaction conditions are the following:—		
45	Water/polymer ratio Carbon/polymer ratio	12 10%	45	
	Hydrolysis temperature Hydrolysis pressure	248°C 42 atmospheres		
50	Residence time in the hydrolyzer	2 hours	50	
	Frequency of carbon discharge every 24 Residence time in the crystallizer Drying temperature	4 hours 80°C		
55	can be carried out in the method of the i obvious that experts in the branch will be consideration, other modifications, witho	odiment has been described and certain changes which nvention under consideration have been outlined, it is a able to make, in view of the description under ut this deviating from the spirit and scope of the	55	
60	invention so that the preceding description	on should be considered only for the purposes of and the scope of this invention should be defined in	60	

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terms of the following clauses:

5	subject to hydrolysis, in the presence of a certain amount of decolorizing carbon, a quantity of polyethylene terephthalate waste with water, at a high pressure and at a temperature which varies between 200°C and 300°C; filter the hydrolyzate also under conditions of high temperature and pressure; crystallize the terephthalic acid of the filtered solution by means of cooling in the crystallization receptor at a final temperature which varies between $-10^{\circ}$ C and	5
	200°C; filter or centrifuge to separate the filtrate crystals; wash and dry the pure terephthalate acid crystals; and, on the other hand, distil the filtrate so as to obtain pure glycol.  2. Continuous method for obtaining pure terephthalic acid and glycol starting from polyethylene terephthalate waste in accordance with what is claimed in clause 1, characterized	
10	furthermore by the fact that the quantity of water used is equal or greater than that required to	10
	keep the terephthalic acid dissolved under hydrolysis conditions.	
	3. Continuous method for obtaining pure terephthalic acid and glycol starting from poly-	
	ethylene terephthalate waste in accordance with what is claimed in clauses 1 and 2, characterized also by the fact that the time of the hydrolysis reaction is between 5 minutes and	
15	6 hours.	15
	4. Continuous method for obtaining pure terephthalic acid and glycol starting from poly-	
	ethylene terephthalate waste in accordance with what is claimed in clauses 1 to 3, also	
	characterized by the fact that the quantity of the decolorizing carbon added is between 0.1%	
20	and 30%, with respect to the quantity of waste treated.  5. Continuous method for obtaining pure terephthalic acid and glycol, starting from	20
20	polyethylene terephthalate waste, in accordance with what is claimed in clauses 1–4, character-	20
	ized also by the fact that the cooling is carried out by venting the pressure in the crystallizer.	
	6. Continuous method for obtaining pure terephthalate acid and glycol, starting from	
25	polyethylene terephthalate, in accordance with what is claimed in clauses 1-5, characterized also by the fact that the final crystallization pressure is equal to atmospheric pressure.	25
25	7. Continuous method for obtaining pure terephathalic acid and glycol, starting from	25
	polyethylene terephthalate waste, in accordance with what is claimed in clauses 1–5, character-	
	ized also by the fact that the final crystallization pressure is lower than atmospheric pressure but	
00	greater than 1 mm Hg absolute, using a vacuum device.	00
30	8. Continuous method for obtaining pure terephthalic acid and glycol, starting from polyethylene terephthalate waste, in accordance with what is claimed in clauses 1–5, also	30
	characterized by the fact that the final crystallization pressure is greater than the atmospheric	
	pressure but less than 15 atmosphere.	
	9. Continuous method for obtaining pure terephthalic acid and glycol, starting from	
35	polyethylene terephthalate waste, in accordance with what is claimed in clauses 1-4, character-	35
	ized also by the fact that the cooling is carried out by means of a cold surface, such as a jacket in the crystallizer, or by means of a coil in the crystallizer, or using both a jacket and a coil.  10. Continuous method for obtaining pure terephthalic acid and glycol, starting from	
	polyethylene terephthalate waste, in accordance with what is claimed in clauses 1-4, also	
40	characterized by the fact that the cooling is carried out by means of any combustion of a cold	40
	surface and a simultaneous venting of the pressure.  11. Continuous method for obtaining pure terephthalic acid and glycol, starting from	
	polyethylene terephthalate waste, in accordance with what is claimed in clauses 1–10, also	
	characterized by the fact that the crystallization time is between 5 minutes and 5 hours.	